Exhibit 5

				Plaintiffe! I	ist of Custodia	ns for ZHP and Its Subsidiaries as of 12/10/2019
				Fiamuns L	ist of Custodia	us for Ziff and its Subsidiaries as of 12/10/2019
First Name	Last name	Title	Department	Co.	Status	Explanation
	1	ı	I	ı	1	Senior Management
		Executive Vice				
		President of ZHP,				
		CEO of Huahai				
		US, Prinston,				
		Solco, and PrinJohnson;				Identified in Prinston's core document production more times than we can reference. Most senior manager involved in the da
		Legal		ZHP, Huahai		to-day aspects of management for ZHP, Prinston, & Huahai US. Fully involved in valsartan contamination issues from initia
		Representative of		US, Prinston,		reporting to FDA to present. Identified as a planned attendee in Prinston's 6.18.18 Email Meeting Request to FDA initially
	_	Shaghai Syncores		Solco, Syncores,		advising the FDA of Valsartan's contamination (PRINSTON00000050). Led ZHP's investigation & response to the FDA's
Jun	Du	and Prinbury		and Prinbury	Disputed	483 observations made during the FDA inspection of Chuannan site 7.23.18 to 8.3.18 (PRINSTON0075644)
		Chairman and		ZHP and		He is the most senior officer in ZHP, and was the chairman of the board of directors. He was involved in a high level into the
Baohua	Chen	General Manager		PrinJohnson	Disputed	valsartan contamination investigation. Jun Du reports directly to him.
		5				ment or "Analysis and Testing Center"
			Analytical			
Min	Li	VP	Operations	ZHP	Agreed	
		m:	at .			Org Chart; involved in testing tetrazole reaction in Valsartan, which is a crucual part of the manufacturing process resulting
Iun	Wana	Finished Dose	Chuannan Branch Analytical Center	ZHP	Dienuted	in the contamination PRINSTON0074146. On 12/4/19, ZHP would not tell Plaintiffs the extent to which this person was involved in Valsartan.
Jun	Wang	Gase Stage Team	Anaiyucai Center	£111	Disputed	III voivou III v aisaitaii.
	Wang or					Is listed as a user on the chromatograms, which can detect nitrosamines, on the December 2013 solvent change submission to
	possibly just					the FDA PRINSTON00073055. As of 12/4/19, ZHP could not identify this person, despite the provision of the referenced
Qing	Wangqing			ZHP	P - 10.8.19	document from Core Discovery.
		Analytical and				
		Testing	CI D 1			Org Chart. Also referenced in the text version of PRINSTON0076941, where he is designated as a member of the Quality
XiaXia	Kuang	Department Manager	Chuannan Branch Analytical Center	7HP	P - 11.22.19	Control Team, which conducted the testing of Valsartan for impurities. As of 12/4/19, ZHP could not identify this person, despite the provision of the referenced document from Core Discovery and the person's presence on its organization chart.
Z KIUZ KIU	ruung	141dHuger	2 marytical Center	ZIII	1 11.22.17	despite the provision of the referenced document from core biscovery and the persons presence on its organization chart.
		Finished Dose				The org chart shows that this person had a significant role in ZHP's Analysis and Testing Department. ZHP has not provided
CI	37	Liquid Phase	Chuannan Branch	ZIID	D 11 22 10	Plaintiffs any additional information on this individual. On 12/4/19, ZHP refused to discuss the extent of this custodian's
Chun	Yang	Team	Analytical Center	ZHP	P - 11.22.19	involvement in Valsartan.
		Raw Material				
C	71	Central Control	Chuannan Branch Analytical Center	ZUD	P - 11.22.19	The org chart shows that this person had a significant role in ZHP's Analysis and Testing Department. ZHP has not provided
Guang	Zheng	Instrument Team	Analytical Center	ZHP	P - 11.22.19	Plaintiffs any additional information on this individual.
		Raw Material				
		Central Control Physico-Chemical	Chuannan Branch			The org chart shows that this person had a significant role in ZHP's Analysis and Testing Department. ZHP has not provided
XianLiang	Zhang	Team		ZHP	P - 11.22.19	Plaintiffs any additional information on this individual.
						Quality
Wai	Chana	VP	API Quality	ZIID	A amaad	
Wei	Cheng	vr	Assurance Quality Assurance	ZHP	Agreed	
			(API Division-			
Jucai	Ge	Director	Chuannan Site)	ZHP	Agreed	
		_	_			
			Quality Assurance			
			Manager (API			
			Division- Chuannan Site,			
Yuelin	Hu	Assistant Director	East Zone)	ZHP	Agreed	
			Corporate Quality			
Baozhen	Chen	Director	Assurance	ZHP	Agreed	
			Quality Assurance at Chuannan Site			
Dongqin	Wang	Assistant Director	East Zone	ZHP	Agreed	
			Finished Dose			
			Quality Assurance			
Min1:	7hor~	Director	and Quality	ZUD	Agreed	
Minli	Zhang	Director	Control	ZHP	Agreed	
			Quality Control (API Division-			
Qiangming	Li	Director	Chuannan Site)	ZHP	Agreed	
	1		,	1		1

Case 1:19-md-02875-RMB-SAK Document 316-5 Filed 12/10/19 Page 3 of 10 PageID: 4910

			API Quality			
Wenbin	Chen	Analysis Director	Research	ZHP	Agreed	
Wenquan	Zhu	Director	API Quality Research	ZHP	Agreed	
			Quality Assurance			Part of core ZHP group identified as meeting attendees regarding the contamination in Duane Morris 6.21.19 correspondence
Xiaohong	Zhao	Director	(Chuannan Site)	ZHP	Disputed	to FDA (PRINSTON00073192)
						Prepared Method Validation Report of Valsartan PRINSTON00031401 (According to published literature, "[f]or the
						pharmaceutical industry, method validation is crucial to ensure the product quality as regards both therapeutic efficacy and patient safety. The most critical step in validating a method is to establish a protocol containing well-defined procedures and
						criteria. A well planned and organized protocol, such as the one proposed in this paper, results in a rapid and concise method
						validation procedure for quantitative high performance liquid chromatography (HPLC) analysis. "See https://conservancy.umn.edu/bitstream/handle/11299/172043/cop_article_496630.pdf?sequence=1&isAllowed=y; Checked
Hongliang	Wang		Quality	ZHP	Disputed	significant tests for Valsartan impurities See, e.g., PRINSTON00023107; reviewed Method Validation Report of Valsartion for correctness and integrity PRINSTON00022665
Honghang	wang		Quarty	ZIII	Disputed	ior concerness and integrity FRASTO-NOOZZZOOS
Cunxiao			Quality Assurance,			Attended close out meeting on 5.19.17 regarding FDA inspection PRINSTON0074158; Approved the Deviation
(Jenson)	Ye	Vice President	Headquarters	ZHP	Disputed	Investigation Reports (root-cause analysis of the contamination) of Valsartan TEA Process PRINSTON0073443, etc.
Caifeng	Zhao	Manager	Quality Assurance	ZHP	Disputed	Principal Author of ZHP's 11.5.18 Deviation Investigation Report (root-cause analysis of the contamination) of Valsartan TEA Process V1, 2, & 3 PRINSTON0073443, 0075797, 0076100
						Reviewed Validation Report of Analytical Method for Azide in Valsartan for correctnesss and integrity PRINSTON00018712 (the removal of azide during the manufacturing process is suspected to contribute to the
						contamination); Reviewed and approved Validation Report of LC-MS Method for NDBA, another nitrosamine, in Losartan
Y.F.	Chen	Senior Supervisor	API Quality Assurance	ZHP	Disputed	Potassium PRINSTON0074642; reviewed and approved the Validation Report of LC-MS Method for NMBA, another nitrosamine, in Irbesartan PRINSTON0074712
Hu	Guangping	Director	Quality Control	ZHP	Disputed	QC Director regarding Valsartan tests related to sodium nitrite, which is suspected to contribute to the contamination PRINSTON00072721
						Referenced in 35 separate documents in core document production, some regarding significant tests for impurities in
Huang	Rui	Director	Quality Control	ZHP	Disputed	Valsartan. See, e.g., PRINSTON00023107
Xie	Chaojun	Analyst	Quality Control	ZHP	Disputed	QC Analyst who conducted tests regarding zinc chloride and DMF, both substances suspected of contributing to the contamination: PRINSTON00072692, 72743, and 00080383
Qiang	Zhou	Analyst	Quality Research	ZHP	Disputed	Drafted Validation Report of GC-MS Method for Detection of NDMA in Valsartan Tablets and Valsartan and Hydrochlorothiazide Tablets PRINSTON00000326, 36625
Shang	Fei	QP	Quality VP	ZHP	Disputed	Reviewed and approved the third version of the Deviation Investigation (root-cause analysis) into the Valsartan TEA process PRINSTON0076100
Tong	Wu	Analyst	API Quality Research	ZHP	Disputed	Drafted Validation Report of GC-MS Method for Detection of NDMA in Valsartan Tablets and Valsartan and Hydrochlorothiazide Tablets PRINSTON00000326, 93
						Reviewed Comparison of Valsartan USP Method and In-house Method for Integrity and Correctness PRINSTON00064389 (This document compares the impurities found in USP standard Valsartan with those found in ZHP's Valsartan); Reviewed
Wenping	Hu	Team Leader	Quality Research	ZHP	Disputed	data in Hydrochlorothiazide Residual Solvent Method Validation Report PRINSTON00064288
						According to the Valsartan DMF, "[s]he is responsible for QA/GMP auditing and the QC/GMP file management for all
						products" PRINSTON00077850; signed Valsartan cGMP Certification PRINSTON00077832. On 12/4/19, ZHP said this person left ZHP in 2008, and ZHP does not have any of their emails. ZHP could not confirm whether ZHP had any other
Jieyun	Wang	Director	Quality Assurance	ZHP	Disputed	electronic or physical documents from this person.
			Formulation			Reviewed and signed off on Elemental Impurity Risk Assessment Report for Valsartan Tablets PRINSTON00036384; ZHP
Xiaoling	Li	Manager	Quality Control	ZHP	Disputed	admits that this person worked on Deviatio Investigation Report and Out-of-Specification Report of Valsartan.
Zheng	Youqing	Deputy Manager	Quality Assurance	ZHP	P - 10.8.19	Executed numerous cGMP certifications, and many of Plaintiffs claims are based on violations of cGMPs re Valsartan. See, e.g. PRINSTON00038190
Ü						
Zhizhang	Ding	Assistant Manager	Formulation Quality Assurance	ZHP	P - 11.12.19	Reviewed and approved Elemental Impurity Risk Assessment Report for Valsartan Tablets PRINSTON00036384; Signed of on correction of the valsartan manufacturing process intended to prevent future contamination PRINSTON00000471
			•			Analyst of Valsartan tests related to sodium nitrite, which is suspected to contribute to the contamination
Xiong	Fei	Analyst	Quality Control	ZHP	P - 10.8.19	PRINSTON00080367; 71698; 72721
1			Quality Control			
Yinhua	Tang	Assistant Director/Manager	(API Division- Chuannan Site)	ZHP	P - 11.12.19	Reviewed and signed off on Deviation Investigation Reports (root-cause analysis of the contaminat) of Valsartan TEA Process PRINSTON0073443, etc.; attended close out meeting on 5.19.17 regarding FDA inspection PRINSTON0074158
	g	_ in cettor, ividinagei	- maniful Site)		- 11.14.1/	2. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2
			Character Co. 17			Drafted Assay Method Validation Report for Valsartan (Supplement) PRINSTON00009948 (An assay method validation is
Yun	Jin	Supervisor	Chuannan Quality Control	ZHP	P - 11.12.19	the process of determining whether the tests, or assays, are accurate and sufficient; it includes chromatograms); draft Chiral Purity Analytical Method Validation Protocol for TDCI Compound PRINSTON00077405

Case 1:19-md-02875-RMB-SAK Document 316-5 Filed 12/10/19 Page 4 of 10 PageID: 4911

Tian	Zhang	Analyst	Chuannan Quality Control	ZHP	P - 11.12.19	Reviewed and signed off on Assay Method Validation Report for Valsartan (Supplement) PRINSTON00009948 (An assay method validation is the process of determining whether the tests, or assays, are accurate and sufficient; it includes chromatograms); reviewed and signed off on Chiral Purity Analytical Method Validation Protocal for TDCI Compound PRINSTON00077405
Xianhua	Zhang	Deputy Director	Chuannan Quality Control	ZHP	P - 11.12.19	Reviewed and signed off on Assay Method Validation Report for Valsartan (Supplement) PRINSTON00009948 (An assay method validation is the process of determining whether the tests, or assays, are accurate and sufficient; it includes chromatograms); reviewed and signed off on Chiral Purity Analytical Method Validation Protocal for TDCI Compound PRINSTON00077405
Yuping	Chen	Deputy Manager	Formulation Quality Control	ZHP	P - 11.12.19	Author of Elemental Impurity Risk Assessment Report for Valsartan Tablets and Valsartan Hyrdochlorathiazide Tablets PRINSTON00036384, PRINSTON00070161; ZHP admits that this person worked on Deviatio Investigation Report and Out of-Specification Report of Valsartan.
Tuping	Chen	Deputy Manager	Quanty Control	ZIII	111.12.17	or openitudin report of variation.
Tan	Xiao	Senior Supervisor	API Quality Research	ZHP	P - 11.12.19	Drafted Validation Report of GC-MS Method for Detection of NDMA in Valsartan and Hydrochlorothiazide Tablets PRINSTON0000326; reviewed and approved Validation report of the LC-MS method for NDBA in Losartan Potassium PRINSTON0074642; reviewed and approved the Validation Report of LC-MS Method for NMBA in Irbesartan PRINSTON0074712
Nan	Tong	Senior Supervisor	API Quality Research	ZHP	P - 11.12.19	Approved the cGMP compliance for its intended use of Report of GC-MS Method for Detection of NDMA in Valsartan and Hydrochlorothiazide Tablets PRINSTON00000326
X. G., possibly Xingge	Liu	Quality Researcher	API Quality Research	ZHP	P - 11.12.19	Drafted Validation report of the LC-MS method for NDBA, a nitrosamine, in Losartan Potassium and Irbesartan PRINSTON0074177. ZHP has not told Plaintiffs that this person was not also involved in detecting nitrosamines in Valsartan.
Y.Y. possibly Yangyang	Не	Quality Researcher	API Quality Research	ZHP	P - 11.12.19	Drafted Validation Report of LC-MS Method for NMBA, a nitrosamine, in Irbesartan and NMSA in Irbesartan PRINSTON0074712, 75156. ZHP has not told Plaintiffs that this person was not also involved in detecting nitrosamines in Valsartan.
Jian	Ye	Deputy Manager, Group I	Quality Control	ZHP	P - 11.22.19	Put into effect the Method Validation, Report for Assay for Valsartan, which is the process of ensuring that the tests were accurate and adequate PRINSTON00070930
Houming	Zhou	General Detection, Technical Deputy Director		ZHP	P - 11.22.19	ZHP's organization chart shows that this person had a signficant role in ensuring the quality of ZHP's API. ZHP has not provided any additional information regarding this person.
Xiao	Yu	Associate Director	API Quality Study/Research Department	ZHP	P - 11.22.19	ZHP's organization chart shows that this person had a signficant role in ensuring the quality of ZHP's API. ZHP has not provided any additional information regarding this person.
Yang	Han	Supervisor	Quality Research	ZHP	P - 11.22.19	ZHP's organization chart shows that this person had a signficant role in ensuring the quality of ZHP's API. ZHP has not provided any additional information regarding this person.
Chaohua	Bao	Supervisor	API QR	ZHP	P - 11.22.19	Prepared Comparison of Valsartan USP Method and In-house Method PRINSTON00064389; put into effect the Method Validation for Valsartan PRINSTON00022665
Jianzhi	Zhao		QA	ZHP	P - 12.2.19	Audit consistency of document and cGMP compliance in many documents See, e.g., throughout PRINSTON00022331-23004; PRINSTON00010056. Many of Plaintiffs' claims are based on violations of cGMPs. Signed cGMP Certification for Valsartan PRINSTON00022284. Many of Plaintiffs' claims are based on violations of
Meng	Zheng	Manager	QA	ZHP	P - 12.2.19	cGMPs.
V	Char		Plant Analytics	7110	P 12 10 10	Worked extensively on Method Validation Report for Valsartan Residual Solvent Method Cross-Validation Report, which included chromatograms, which can detect nitrosamines PRINSTON00064537. Plaintiffs believe solvent played a critical role in the contamination. ZHP admits that this person worked on Deviatio Investigation Report and Out-of-Specification
Yong	Chen	Material Team	Center	ZHP	P - 12.10.19	Report of Valsartan.
Shuli	Du		Finished Dose Plant Analytics Center	ZHP	P - 12.10.19	Reports to Chen Yuping, who ZHP admits worked on Deviation and Out-of-Specitivation reports related to Valsartan. ZHP has not been able to provide any additional information.
Danyara	Zhang	Sterile Material	Finished Dose Plant Analytics	7ИР	P - 12.10.19	Reports to Chen Yuping, who ZHP admits worked on Deviation and Out-of-Specitivation reports related to Valsartan. ZHP
Danyang	Zhang	Team	Center	ZHP	r - 12.10.19	has not been able to provide any additional information.
Na	Li	Senior Supervisor of Material Review Team	Finished Dose Plant Analytics Center	ZHP	P - 12.10.19	Reports to Chen Yuping, who ZHP admits worked on Deviation and Out-of-Specitivation reports related to Valsartan. ZHP has not been able to provide any additional information.

		Director of Analytics	Finished Dose Plant Analytics			Supervises Chen Yuping, who ZHP admits worked on Deviation and Out-of-Specitivation reports related to Valsartan. ZHP
Pang	Hu	Integrated Room	Center	ZHP	P - 12.10.19	has not been able to provide any additional information.
8						
		Technical				
			Finished Dose			
	71	Quality Research		ZUD	D 12 10 10	ZHP has not been able to provide any additional information about this person, but their title and place in the organization
Fuyu	Zhang	Team	Center	ZHP	P - 12.10.19	chart show that they very likely worked on Valsartan and would have relevant, probative documents in this case.
			Finished Dose			ZHP has not been able to provide any additional information about this person, but their title and place in the organization
		Senior Supervisor				chart show that they very likely worked on Valsartan and would have relevant, probative documents in this case, especially
Ling	Lin	of Stability Team	Center	ZHP	P - 12.10.19	because stability testing could have provided notice of the nitrosamine contamination.
		Deputy Director of				
			Finished Dose Plant Analytics			ZHP has not been able to provide any additional information about this person, but their title and place in the organization
Zeqi	Min	Room	Center	ZHP	P - 12.10.19	chart show that they very likely worked on Valsartan and would have relevant, probative documents in this case.
1			Finished Dose			ZHP has not been able to provide any additional information about this person, but their title and place in the organization
			Plant Quality			chart show that they very likely worked on Valsartan and would have relevant, probative documents in this case, especially
Kaiwei	Zhu	Deputy Manager of Compliance	Management Department	ZHP	P - 12.10.19	because of the person's presence in ~32 Valsartan Batch Records, which document the entire manufacturing process of Valsartan.
Karwer	Znu	от сопірпанес	Department	ZIII	1 - 12.10.17	Y aisartair.
			Finished Dose			
			Plant Quality			
		Validation	Management			ZHP has not been able to provide any additional information about this person, but their title and place in the organization
Yang	Yang	Manager	Department	ZHP	P - 12.10.19	chart show that they very likely worked on Valsartan and would have relevant, probative documents in this case. Manufacturing
		I	Technical for	Ī		Manufacturing
Xiaoming	Liu	Head	Finished Dose	ZHP	Agreed	
			Technical			
Peng	Dong	Deputy Director	Department	ZHP	Agreed	
			ZHP Facility			
			Director API			
			Manufacturing			
Peng	Wang	Senior Director	Chuannan Site, West Zone	ZHP	Agreed	
reng	wang	Schiol Birector		ZIII	Agreed	
			Technical, API			
			Chuannan site West Zone			
			(counterpart to			
Wenling	Zhang	Director Assistant	Peng Dong)	ZHP	Agreed	
			Production			
Wei	Chen	Deputy Director	Department	ZHP	Agreed	
L	L.	Factor Assistant				Identified in Valsartan DMF as a "professional engineer," and "has a wide experience at production technology management
Yongjun	Jin	Director	Manufacturing	ZHP	Disputed	and is responsible for the technological development of all bulk drugs" PRINSTON00077836
		Deputy Director of				
Vanhu	Wana	Production and		ZIID	Diamuta 4	Identified in Valsartan DMF as "responsible for Valsartan" PRINSTON00077836; appears in the text of later documents,
Youhu	Wang	Operation		ZHP	Disputed	such as ZHP's Response to the FDA's August 2018 inspection PRINSTON0074892
		API and				
		Intermediate Plant				
Jiangzhong	Yu	Manager	Manufacturing	ZHP	Disputed	Identified in Valsartan DMF as responsible for all bulk drugs (including intermediates) at ZHP PRINSTON00077836
			API Operations/Facility			
			Operations/Facility Director, API			
			Chuannan Site,			
Lijin	Jiang	VP	East Zone	ZHP	Disputed	Attended close out meeting on 5.19.17 regarding FDA inspection PRINSTON0074158
,						
,			Engineering and			
			Maintenance, API			
Zhengjun	Jia	Deputy Plant		ZHP	Disputed	Attended close out meeting on 5.19.17 regarding FDA inspection PRINSTON0074158

		1	1	1	1	
			Parisaraisa and			
			Engineering and Maintenance, API			
			Chuannan site			
Junhui	Zuo	Director	West Zone	ZHP	Disputed	Attended close out meeting on 5.19.17 regarding FDA inspection PRINSTON0074158
Chunmin	Xu	Vice President		ZHP	Disputed	Identified as Valsartan personnel in DMF PRINSTON00070584, 79392. On 12/4/19, ZHP said this person left ZHP in 2015, and ZHP does not have any of their emails. ZHP could not confirm whether ZHP had any other electronic or physical documents from this person.
Chamin		Vice i resident			Disputed	accumumo nom uno personi
Ruqi	Yao	Factory Assistant Director		ZHP	Disputed	Identified as Valsartan personnel in DMF PRINSTON00070587, 79392. On 12/4/19, ZHP said this person left ZHP in 2012, and ZHP does not have any of their emails. ZHP could not confirm whether ZHP had any other electronic or physical documents from this person.
**		Technical				Identified as Valsartan personnel in DMF PRINSTON00070587, 79392. On 12/4/19, ZHP said this person left ZHP in 2014, and ZHP does not have any of their emails. ZHP could not confirm whether ZHP had any other electronic or physical
Kai	Yang	Manager		ZHP	Disputed	documents from this person.
Meng	Yanhua	General Manager		ZHP	Disputed	The head of ZHP's manufacturing department at the facility that manufactured Valsartan, presumably communicated between her subordinates, other departments, and senior management regarding Valsartan manufacturing.
Lina	Wu	Senior Supervisor	Formulation Technology at ZHP	ZHP	P - 11.12.19	Author of Elemental Impurity Risk Assessment Report for Valsartan Tablets and Valsartan and Hydrochlorothiazide Tablets PRINSTON00036384, 70161
Hu	Zhou	Factory Director		ZHP	P - 11.22.19	Identified as Valsartan personnel responsible for production and technology in DMF PRINSTON00070586
Yan	Fengfeng	Director of Tech	Chuannan Product Technical Center	ZHP	P - 11.22.19	As the second in command at ZHP's Technical Department in the facility manufacturing Valsartan API, this person was responsible for developing the manufacturing process for Valsartan.
Liang	Zunjun	Associate Director	Chuannan Product	ZHP	P - 11.22.19	The assistant to Dong Peng, who ZHP admits is a proper custodian. ZHP has not given Plaintiffs any additional information regarding this person.
Liang	Zunjun	Associate Director	reclinical Center	ZIII	1 - 11.22.19	regarding this person.
			Chuannan Product			Manager in the department that developed the manufacturing process for Valsartan. ZHP has not given Plaintiffs any
Hu	Zhichen	Manager	Technical Center	ZHP	P - 11.22.19	additional information regarding this person.
			Chuannan Product			This person's title is Valsartan. They are in the department that developed the manufacturing process for Valsartan. ZHP has
Zhan	Xiaohui	"(Valsartan)"	Technical Center	ZHP	P - 11.22.19	not given Plaintiffs any additional information regarding this person.
		Deputy Head of	Finished Dose			Reviewed and signed off on numerous Batch Reocrds, which document then tire manufacturing process See, ie.,
		Technology	Plant Technology			PRINSTON00068034; also supervised Lina Wu, who authored Author of Elemental Impurity Risk Assessment Report for
Lingfang	Wang	Guarantee Unit	Center	ZHP	P - 12.10.19	Valsartan Tablets and Valsartan and Hydrochlorothiazide Tablets PRINSTON00036384, 70161
				7110	D 121010	Wild Miles and April 1997 April 1
Hui	Wang	Director	Quality Control	ZHP	P - 12.10.19	Worked on Valsartan's solvent testing that likely contributed to contamination PRINSTON00078435
			Finished Dose			
lie	Cheng		Plant Technology Center	ZHP	P - 12.10.19	ZHP has not been able to provide any additional information about this person, but their title and place in the organization chart show that they very likely worked on Valsartan and would have relevant, probative documents in this case.
Jie	Cheng	1	Center	LIII	1 - 12.10.19	tenart show that they very ricery worked on varisarian and would have relevant, probative documents in this case.
						ZHP has not been able to provide any additional information about this person, but their title and place in the organization
			Finished Dose Non			chart show that they very likely worked on Valsartan and would have relevant, probative documents in this case, especially
Juan	Tao	Manager of Production	Sterile Production Division	ZHP	P - 12.10.19	because they worked on numerous Batch Records for Valsartan, which document the manufacturing process, including its testing.
- auii	1100		2.1101011		12.10.17	Regulatory
					1	
					1	
Lihong (Linda)	Lin	Director	Regulatory Affairs Department	ZHP	Agreed	
(Dindu)	2.11	Z ii cettoi	2 oparament		51000	
					1	
Yanfeng	Lin	Deputy Director /	Danulet + 00 '	ZUD	A ana - 4	
(Lucy)	Liu	Manager	Regulatory Affairs	ZHP	Agreed	
Ying	Xiong	Supervisor	Regulatory Affairs	ZHP	Agreed	
Ting (Ada)	Zhou	Assistant Director	Regulatory Affairs	ZHP	Agreed	
- ` /	•	•				

Case 1:19-md-02875-RMB-SAK Document 316-5 Filed 12/10/19 Page 7 of 10 PageID: 4914

				T		,
			API Patent,			Head of patents at ZHP, which has patents regarding Valsartan, one of which is for the manufacturing process that led to the
Jie	Lin	Deputy Manager	Regulation Affairs	ZHP	Disputed	contamination.
			API R&D,			Manager of API Research and Development in the Regulatory Department that handled Valsartan. ZHP has not given
Sun	Dengxue	Manager	Regulation Affairs	ZHP	P - 11.22.19	Plaintiffs any additional information regarding this person. Procurement
Min	Hu		Purchasing	ZHP	Agreed	11 ocurement
Weiwei	Xu Xu		Purchasing Purchasing	ZHP ZHP	Agreed	
Min Zheng	Gaozhe		Purchasing	ZHP	Agreed Agreed	
				ı	ı	API Sales
Minda	Cai	Former VP	API Sales	ZHP	Agreed	
			Business			
Jie	Wang	VP	Development	ZHP	Agreed	
Sheng	Zhong	Director	Business Development	ZHP	Agreed	
Sheng	Linong		Business		rigirea	
Hongchao	Li	Manager	Development Business	ZHP	Agreed	
			Development, API			
Mi	Xu	ļ	Sales	ZHP	Agreed	
						Co-head of ZHP's Sales Department, which handled Valsartan. ZHP has said that she is a recent hire, but has not said how
Yihua	Wang	Vice President	Sales	ZHP	Disputed	recent. ZHP also admits that she has knowledge of the Valsartan recall.
Eamariana	Tono	Manager Assistant, Business Group I	Calas	ZHP	Diamortad	Manager in the Sales Department who handled Key Accounts for Valsartan. ZHP has not told Plaintiffs that Fengyang Tang and Lina Wang did not both work on Valsartan.
Fengyang	Tang	Business Group I	Sales	ΖПΓ	Disputed	and Lina wang did not both work on varsartan.
		Manager Assistant,				Manager in the Sales Department who handled Key Accounts for Valsartan. ZHP has not told Plaintiffs that Fengyang Tang
Lina	Wang	Business Group II Director of	Sales	ZHP	Disputed	and Lina Wang did not both work on Valsartan.
		Customer Service				As the Director of Customer Service, this person very likely dealt with customer complaints regarding Valsartan. ZHP has no
Tong	Zengyuan	Department	Sales	ZHP	P - 11.22.19	provided any additional information regarding this person.
		Manager of				As the manager of the Document Group in the Customer Service Department, this person is a critical cutodian. ZHP has not
Wu	Yuehua		Sales	ZHP	P - 11.25.19	given Plaintiffs any additional information regarding this custodian.
		Assistant Manager				The Aris of the Court of the Co
Wang	Haigun	of Office Work Group	Sales	ZHP	P - 11.25.19	The Assistant Manager in the Customer Service Department. ZHP has not given Plaintiffs any additional information regarding this custodian.
			Ī		1	Hauhai US
		Executive Vice		Huahai US		
		President and		(linkedin),		
		Chief Scientific Officer for		Prinston (website), and		
N. 1.	6	Prinston & Huahai		Solco (affidavits	<u>,</u>	
Xiaodi	Guo	US		of service)	Agreed	
		1		Vice President at		
				Huahai US since		
		1		2005 (Linkedin), Sr. Vice		
				President at		
		1		Prinston		
				(Linkedin), & President at		
** .	***	President since		Solco (Prinston's	<u>,</u>	
Hai	Wang	2005		website)	Agreed	
				**	<u> </u>	
John	Iozzia	Director	Sales	Huahai US	Agreed	
71 .	GI.	1		11 1 : 12	D:	The transfer of the transfer o
Zhi	Chen			Huahai US	Disputed	Listed as contact for ZHP's US Agent to the FDA for Valsartan, Huahai US PRINSTON00072295
		<u></u>		**		Author of Cover Letter re: Valsartan ANDA PRINSTON00066953-54; Listed as contact for the US Agent, Huahai US, for
Qun (Kathy)	Zhang	Senior Manager	Regulatory Affairs	Huahai US	Disputed	Valsartan PRINSTON00073120; Same PRINSTON00072212

			T			
Min (Michelle)	Hu	Manager of Business Development and Senior Director of Business Development		Huahai US (2007-2012) and Prinston (2012- Present), respectively	Disputed	Given her title and Huahai US's and Prinston's significant role in bringing ZHP's Valsartan to the United States, Plaintiff's believe that she has important, relevant, and probative documents in this case. The ZHP defendants do not dispute this. They simply say that other custodians "have superior information related to Valsartan."
Yiming	Tang				P - 11.22.19	Contact Person for ZHP's U.S. Agent to FDA for Valsartan PRINSTON0076915 and many other documents regarding Huahai US as ZHP's U.S. agent to the FDA for Valsartan PRINSTON00010626, PRINSTON00008984, PRINSTON00078888, etc.
					l	Prinston
Xiaodi	Guo	Executive Vice President and Chief Scientific Officer for Prinston & Huahai US		Huahai US (linkedin), Prinston (website), and Solco (affidavits of service)	Agreed	
Hai	Wang			Vice President at Huahai US since 2005 (Linkedin), Sr. Vice President at Prinston (Linkedin), & President at Solco (Prinston's website)	Agreed	
Chris	Keith	Senior Vice President / VP	Sales	Solco/Prinston at some point as VP	Agreed	
Remonda	Gergis	VP	Quality Assurance	Deinston	Agreed	
	Wang	VP	Head of Regulatory Affairs		Agreed	
Minfa	Wang	VP	Analytical Operations and Quality Control	Prinston	Disputed	She is a senior officer of Prinston, in charge of Analytical Operations and Quality Control. Identified as a planned attendee in Prinston's 6.18.18 Email Meeting Request to FDA initially advising the FDA of Valsartan's contamination (PRINSTON00000050).
Wei (Helena)	Tong	Director	Regulatory Affairs	Prinston	Disputed	Listed on ANDA Documents, Signed ANDA, and routinely corresponded with the FDA. For example, PRINSTON00037372, PRINSTON00054685, , PRINSTON00031790. Her full name appears in 106 documents from Core Discovery.
Guangyu	Luo	Associate (2013- 2015), Senior Associate (2015- 2016), Specialist (2016-2017), Manager (2017- Present)		Prinston	Disputed	Signed ANDA for Valsartan PRINSTON00066560, PRINSTON00033715, PRINSTON00034648, PRINSTON00066787, PRINSTON00066707, PRINSTON00066848; "Regulatory Affairs, mainly on regulatory submissions (from draft to e-CTD compilation), pharmacovigilance (AE handling and submission) and generic labeling (content, artwork and SPL)." -Linkedin
Lesley	Zhu	SVP, Business Development & Portfolio	Sales	Prinston	Disputed	Received email from Frederick Ball re: FDA correspondence along with Jun Du; Same PRINSTON0073198; Cc'd on letter to FDA with Jun Du PRINSTON00079000; Sent letter to FDA re: authorizing the FDA to share information with the EMA PRINSTON0070470-71; PRINSTON0073192-96; Cc'd on letter to FDA with Jun Du PRINSTON00079001-02
Min (Michelle)	Hu	Manager of Business Development and Senior Director of Business Development		Huahai US (2007-2012) and Prinston (2012- Present), respectively	Disputed	Given her title and Huahai US's and Prinston's significant role in bringing ZHP's Valsartan to the United States, Plaintiffs believe that she has important, relevant, and probative documents in this case. The ZHP defendants do not dispute this. They simply say that other custodians "have superior information related to Valsartan."

Case 1:19-md-02875-RMB-SAK Document 316-5 Filed 12/10/19 Page 9 of 10 PageID: 4916

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Peng	Shang	Senior Associate of Business Development/Sup ply Chain (2016- Present)		Prinston	Disputed	Given this person's title and Prinston's significant role in bringing ZHP's Valsartan to the United States, Plaintiffs believe that she has important, relevant, and probative documents in this case. The ZHP defendants do not dispute this. They simply say that other custodians "have superior information related to Valsartan."
Nina	Zhang	Specialist	Regulatory Affairs	Princton	P - 12.2.19	Regulatory Affairs Approval for Corrected Valsartan Manufacturing Process PRINSTON00000479
Ning	Zhang	Specialist	Regulatory Arrairs	FIIIISIOII	r - 12.2.19	Solco
Xiaodi	Guo	Executive Vice President and Chief Scientific Officer for Prinston & Huahai US		Huahai US (linkedin), Prinston (website), and Solco (affidavits of service)	Agreed	
				Vice President at Huahai US since 2005 (Linkedin), Sr. Vice President at Prinston (Linkedin), & President at Solco (Prinston's		
Hai	Wang			website)	Agreed	
		Senior Vice		Solco/Prinston at some point as		
Chris	Keith	President / VP	Sales	VP	Agreed	
David	Aures	Vice President (2018-Present); Executive Director, Sales (2016-2018); Senior Director of National Accounts (2014-2016); Director of National Accounts (2012-2013); National Account Manager (2010-2012)		Salca	Agreed	
David	Ayres	2012)	Sales	Solco	Agreed	
Matthew	Arnold	National Account Manager from 2016-2019		Solco	Disputed Shangha	Author of Only Solco-stamped document, a list of Valsartan customers. He must have have many other documents regarding these customers. Solco has not said otherwise. ii Syncores Technologies, Inc.
				Shanghai		
Eric	Gu	Manager		Syncores	Agreed	
An	Jianguo	Vice President	Process Research and Development	Shanghai Syncores	Agreed	
			Process Research	Shanghai		
Huang	Luning	Vice President	and Development	Syncores	Agreed Pi	rinbury BioPharm Ltd.
Eric Wen- Chien	Tsai	General Manager		Prinbury	Agreed	
Tiyun (Tina)	Liu	Senior Scientist		Prinbury	Agreed	
	Tian	Senior Director		Prinbury	Agreed	
Taufaung (Alex)	Liu	Scientist		Prinbury	Agreed	

Case 1:19-md-02875-RMB-SAK Document 316-5 Filed 12/10/19 Page 10 of 10 PageID: 4917

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Dachuan	Zhao	Executive Deputy General Manager of Prinbury		Prinbury	P - 11.12.19	Attended close out meeting on 5.19.17 regarding FDA inspection, identified as Vice President of Analytical Shanghai R&D Center PRINSTON0074158; "Dr. Zhao has comprehensive and in-depth understanding and rich experience in key areas such as product development (QbD), analysis, product declaration, quality control, quality assurance, FDA regulations, and FDA audits."-Prinbury website (via Google Translate). Identified in core discovery as Vice President. Analytical, Shanghai R&D Center PRINSTON0076915. Prinbury developed the finished dose manufacturing process for Valsartan, including many of the tests conducted during that process, as shown below.
Li or Lily	Tan	Director	Project Management	Prinbury	P - 11.12.19	Worked on numerous important Valsartan documents, many of which involed chromatography, which can detect nitrosamines: Reviewed and Signed Off on Method Robustness Study Report re: Content Uniformity Determination of Valsartan and Hyrdochlorothiazide PRINSTON00039541; Reviewed and Signed Off on Method Validation Report on Identification, Assay and Content Uniformity Determination of Valsartan PRINSTON00020805; Approved Method Verification Report for Valsartan PRINSTON00039715; Reviewed and signed off on Dissolution Comparison Study Report of Valsartan PRINSTON00029309; Approved Method Validation Report re: Dissolution Test of Valsartan and Hyrdochlorothiazide PRINSTON 00039663; Approved Method Verification Report re: Identification and Assay Determination of Valsartan and Hyrdochlorothiazide PRINSTON00039439; Approved Method Verification Report re: Organic Impurities Determination for Valsartan and Hyrdochlorothiazide PRINSTON00039499; Approved Submission Batch Stability Study Report for Valsartan and Hyrdochlorothiazide PRINSTON00039499; Approved Submission Batch Stability Study Report for Valsartan and Hyrdochlorothiazide PRINSTON00064193; Reviewed and Signed Off on Method Robustness Study Report re: Dissolution Determination of Valsartan and Hyrdochlorothiazide PRINSTON00039645; Approved Method Verification Report re: Dissolution Test of Valsartan and Hyrdochlorothiazide PRINSTON00039800; "12 years of experience in drug development, analysis, quality management and project management. Good at the establishment and management of quality system and project managemen system. Familiar with US FDA policies, regulations, cGMP specifications and ICH guidelines, and proficient in FDA registration requirements. Participated and successfully passed GMP audits of FDA, EMEA and CFDA." -Prinbury website (via Google Translate).
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Wei	Tian	Deputy General Manager of Analysis and Research		Prinbury	P - 11.12.19	This person was previously on the Prinbury website, but no longer is. Wei Tian's title and Prinbury's work on Valsartan establish that Wei Tian is likely to have worked on Valsartan related issues and potentially had involvement in and/or is knowledgeable as to how it was tested, among other things.
						Worked on numerous important Valsartan documents, many of which involed chromatography, which can detect nitrosamines: Prepared Method Verification Report re: Identification and Assay Determination of Valsartan and Hyrdochlorothiazide PRINSTON00039439; Prepared Method Verification Report re: Organic Impurities Determination for Valsartan and Hyrdochlorothiazide PRINSTON00039715; Prepared Submission Batch Stability Study Report re: Valsartan and Hyrdochlorothiazide PRINSTON00064193; Prepared USP Method Equivalence Study Report re: Content Uniformity Determination of Valsartan and Hyrdochlorothiazide PRINSTON00039499; Prepared Method Validation Report re: Content Uniformity Determination of Valsartan and Hyrdochlorothiazide PRINSTON00039560; Prepared Dissolution Comparison Study Report re: Valsartan PRINSTON00029309; Prepared USP Method Equivalence Study Report re: Dissolution Test of Valsartan and Hyrdochlorothiazide PRINSTON00039603; Prepared Method Robustness Study Report re: Dissolution Determination for Valsartan and Hyrdochlorothiazide PRINSTON00039645
Shine	Chen	Scientist		Prinbury	P - 11.12.19	
Ashley	Lin	Associate Scientist		Prinbury	P - 11.12.19	Prepared Method Robustness Study Report for Content Uniformity Determination, using Chromatography, which can detect nitrosamines, of Valsartan PRINSTON00039541; Prepared USP Method Equivalence Study Report, using Chromatography, which can detect nitrosamines, re: Dissolution Test of Valartan PRINSTON00021157; Prepared Method Validation Report for Dissolution Test, using Chromatography, which can detect nitrosamines, of Valsartan and Hydrochlorothiazide PRINSTON00039663
Yulu (Luke)	Wang	Executive Director		Prinbury	P - 11.12.19	Reviewed and approved many Master Production Batch Record Approval See, e.g., PRINSTON00064070, PRINSTON00063906, PRINSTON00063742, PRINSTON00038525. Batch Records contain actual data and step by step process for manufacturing each batch. Batch manufacturing record is like a proof that batches were properly made and checked by quality control personnel. See https://process-xe.sarjen.com/batch-manufacturing-records-important-according-gmp/.
		G : .:.		D: 1	D 12210	Prepared Method Validation Report on Identification, Assay and Content Uniformity Determination of Valsartan, using
Angela	An	Scientist		Prinbury	P - 12.2.19	Chromatography, which can detect nitrosamines PRINSTON00020805